



SmartPA Criteria Proposal

| Drug/Drug Class: | Selzentry Clinical Edit | | | |
|----------------------------|-----------------------------------------------------------------|--|--|--|
| First Implementation Date: | April 7, 2010 | | | |
| Proposed Date: | December 17, 2020 | | | |
| Prepared for: | MO HealthNet | | | |
| Prepared by: | MO HealthNet/Conduent | | | |
| Criteria Status: | ⊠Existing Criteria □Revision of Existing Criteria □New Criteria | | | |

Executive Summary

Purpose: Ensure appropriate utilization and control of Selzentry® (maraviroc)

Why Issue Selected: Selzentry® (maraviroc) is a CCR5 (C-C chemokine receptor type 5) co-receptor antagonist indicated for the treatment of only CCR5-tropic HIV-1 infection with other antiretroviral agents. In 2016, Selzentry received FDA approval to expand its indication to include treatment of pediatric patients 2 years of age and older. Selzentry works by selectively binding to the human chemokine receptor CCR5 present on the cell membrane and preventing the interaction of HIV-1 gp 120 and CCR5; this interaction is necessary for CCR5-tropic HIV-1 to enter cells. Selzentry is not recommended in patients with CXCR4tropic or dual/mixed HIV-1 as it is ineffective in these cases. The Trofile® test is used to determine if a patient's virus is suitable for a CCR5 co-receptor antagonist therapy; the results of the Trofile test will determine if Selzentry is an appropriate treatment choice for the patient. Also, as compared to treatment with Sustiva® (efavirenz), treatment-naïve adults treated with Selzentry experienced more virologic failure and lamivudine resistance; therefore, Selzentry is not recommended for use in treatment-naïve patients. Due to the specific approved indication, MO HealthNet will impose clinical criteria to ensure appropriate utilization of Selzentry.

Program-Specific Information:

| ; | Date Range FFS 10-01-2019 to 9-30-2020 | | | | | | | |
|---|----------------------------------------|--------|-------------|-------------------------|--|--|--|--|
| | Drug | Claims | Spend | Average Spend per Claim | | | | |
| | SELZENTRY 25 MG TABLET | 0 | • | - | | | | |
| | SELZENTRY 75 MG TABLET | 0 | - | - | | | | |
| | SELZENTRY 150 MG TABLET | 22 | \$32,009.46 | \$1,454.97 | | | | |
| | SELZENTRY 300 MG TABLET | 43 | \$69,570.80 | \$1,617.92 | | | | |
| | SELZENTRY 20 MG/ML SOLN | 0 | - | - | | | | |

| Type of Criteria: | ☐ Increased risk of ADE☒ Appropriate Indications | □ Preferred Drug List☑ Clinical Edit |
|-------------------|-----------------------------------------------------------------------------|-----------------------------------------------------------------|
| Data Sources: | ☐ Only Administrative Databases | □ Databases + Prescriber-Supplied |

Setting & Population

- Drug class for review: Selzentry® (maraviroc)
- Age range: All appropriate MO HealthNet participants aged 2 years and older

Approval Criteria

- Participant is aged ≥ 2 years AND
- Participant is HIV infected AND
- Participant has history of positive viral tropism for CCR5-tropic HIV AND
- Participant is not treatment naïve (participant has been on antiretroviral medication before) AND
- Participant is currently on additional antiretroviral medication besides Selzentry (maraviroc)

Denial Criteria

Therapy will be denied if all approval criteria are not met

| Required Documentation | | | | | | | | |
|---------------------------------------|---|---------------------------|---|--|--|--|--|--|
| Laboratory Results: MedWatch Form: | X | Progress Notes: Other: | X | | | | | |

Disposition of Edit

Denial: Exception code "0682" (Clinical Edit)

Rule Type: CE

Default Approval Period

1 year

References

- Selzentry (maraviroc) [package insert]. Research Triangle Park, NC: ViiV Healthcare; July 2018.
- Panel on Antiretroviral Guidelines for Adults and Adolescents. Guidelines for the Use of Antiretroviral Agents in Adults and Adolescents with HIV. Department of Health and Human Services. https://clinicalinfo.hiv.gov/sites/default/files/guidelines/documents/AdultandAdolescentGL.pdf. Accessed October 28, 2020.